### DECLARATION UNDER 37 C.F.R. § 1.132

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۱	First Inventor	NORTON
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	Examiner	King, Felicia C.
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Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

SIR:

- I, Mark Norton, declare that:
- I am one of the co-inventors of the present application. I have been involved in all stages of the development of the subject matter disclosed in the present application.

  I am also aware of the pending claims in the present application, as well as the Office Action mailed December 19, 2008.

  | Property of the present application | Property of the property of the present application | Property of the present application | Property of the pro
- 2. The present invention is directed to a novel discovery by the inventors, namely that consumer liking, i.e. taste preference, for a coffee product can be affected by the addition of a single naturally occurring component in coffee, linalool. The principal consumer sensory characteristics of coffee are its roast quality/bitterness and acidity, as determined by consumer sensory evaluations of numerous diverse coffee types. An extensive independent study was conducted and published in an article by the European Sensory Network Food Research Association, entitled "European Sensory and Consumer Study—A Case Study on Coffee" by J.A. McEwan (relevant excerpts attached in Appendix A). In the study, other attributes were described which play a minor role in characterizing the coffee flavor. Inherent flavors in coffee are described as grassy, green,

citrus, floral, fruity, cereal, roasted, caramel, bitter, woody and winey, as well as other attributes.

- 3. The flavor of coffee is the result of the coffee beans and the roasting conditions used to produce the roast coffee blend. Depending on the beans and roasting conditions, the coffee blend will be classified by the origin of the coffee bean, e.g., Sumatra, Columbia, Kona, or by roasting conditions, e.g., French roast or Espresso. Although there will be variations in the taste of these blends, collectively they would all be referred to as non-flavored or natural coffee.
- 4. Coffee itself is a complex product which includes many compounds. Of these flavor compounds, only a few have been previously considered relevant to the taste and aroma in the final coffee beverage. Of the previously considered relevant compounds, they can be classified into groups by the flavor attribute with which they are associated when present within coffee beverages. One example of a flavor group is roasted-nutty.
- 5. While coffee has natural attributes, in recent years, coffee manufacturers have added an additional flavor to the coffee product, i.e. a non-coffee flavor, which includes hazelnut, chocolate, vanilla, raspberry, Irish crème, toffee, orange, amaretto and marshmallow. The resulting product is referred to as "flavored coffee" to distinguish it from natural coffee product.
- 6. When processing coffee beans with the intent of positively affecting the intensity of one flavor attribute, commonly there is a negative affect on at least one other flavor attribute. This can be linked to the chemistry of coffee flavor compounds which undergo chemical reactions depending on the roast conditions. Furthermore, due to

different chemical reactions and the roast conditions, it is unpredictable as to what the effect altering even a single flavor, such as adding a flavorant, will have in the final coffee taste.

- 7. As noted in paragraph 2 above, one naturally occurring component in brewed coffee is linalool. However, prior to the present invention, linalool, was not believed to be of any "major" importance to coffee flavor and/or consumer liking. See, e.g., Blank-I; Sen-A; Grosch-W, "Potent odorants of the roasted powder and brew of Arabica coffee," Zeitschrift-fuer-Lebensmittel-Untersuchung-und-Forschung; 195 (3) 239-245, 1992 (hereinafter "Blank et al."), attached to this declaration as Appendix B, which assesses 3,7-dimethylocta-1,6-dien-3-ol, i.e. linalool, as being unimportant in the overall contribution to roasted coffee flavor. The Blank et al. study included 38 compounds that the researchers could detect with their equipment. This included linalcol. The summary on the front page highlights the 13 compounds that the group found to be important to the overall perception of coffee - linalool is absent. Further, Ivor Flament, "Coffee Flavour Chemistry," Wiley Press, pp. 104-105 (hereinafter "Flament," attached in Appendix C) suggests it is probably unimportant at the concentrations typically found in brewed coffee. Still, another important reference makes no mention of it within a list of important compounds, namely Illy, A; Viani, R; "Espresso Coffee: The Science of Quality," Elsevier Academic Press, p. 201 (hereinafter "Illy", attached in Appendix D).
- 8. Linalool is typically present in roast and ground coffee product in an amount between 30 μg/kg and 4700 μg/kg, and in brewed coffee in an amount of 1 μg/l to about 30 μg/l. Linalool, in isolation, is characterized as having a fruity-floral character (see, e.g., Flament, Appendix C).

- 9. Prior to the present invention, linalool was considered to be an undesirable component in coffee (Flament, pp. 104-15, Appendix C). Specifically, presence of linalool was associated with providing an "undesirable note in disharmony with the notes of a roasted coffee." (*Id.*) Linalool is further described as being a "potent odorant [] of roasted powder of Arabica coffee but not in brew." (*Id.*, citing Blank et al.) Further, prior to the present invention, one would not have believed that it would be desirable to add a fruity-floral character in a natural coffee flavor product. Therefore, one did not previously add additional linalool to a coffee product (Flament, Appendix C).
- 10. Discovery of the present invention proceeded by first taking coffee compounds and identifying their respective contributions to specific sensory coffee attributes in order to assess the importance that each attribute contributes to consumer liking. Prior to the present invention, we were not aware of any other study which determined in detail which groups of compounds were responsible for specific attributes contributing to consumer liking. During our study, we determined how to enhance, degrade or decouple several flavor and aroma attributes by the addition of associated groups of coffee flavor compounds within a robust statistical design. This method permitted us to assess consumer reaction towards enhanced levels of each individual attribute and did not generate unfamiliar attributes within the context of pure coffee. As a result, we were able to "decouple" coffee attributes in order to accurately assess which flavor and aroma attributes contributed to consumer liking. We then determined consumer responses to flavor attributes, i.e. components associated with driving coffee product liking. In addition, we evaluated consumer responses for components that previously had

only been associated with contributing to subtleties of the coffee product profile on a component-by-component basis.

- 11. Through our experimentation, we surprisingly determined that a single chemical, linalool, which previously was not known to be of primary importance to affect or be associated with sensory attributes, let alone to be important at all in the overall coffee flavor, drove significant consumer liking. Further, it was determined that enhanced consumer liking was attributed to linalool in a roast and ground coffee having levels of linalool, as measured in the roast and ground product, of at least 6000 µg/kg using the Likens method. Based on these results, the amount of linalool in roast and ground coffee, one can extrapolate the data to soluble coffee having levels of at least 2000 µg/kg of soluble coffee solids present in the soluble coffee.
- 12. Some coffee experts come to regard the delicate Arabica coffee from East Africa as high quality, e.g., Ethopian Sidamo and Ethiopian Djimmah. These origins of coffee are relatively rare and expensive. We have identified that a key differentiating chemical signature within this family of coffee is the higher level of linalool, and found through the course of this work that elevating the level of linalool in more common, lower-linalool variants that we can move the flavor character towards the more expensive and rare Ethiopian coffee.
- 13. Further, surprisingly we determined that fruity and floral attributes substantially drove consumer liking. Prior to the present determination, it was unknown in the art that fruity and floral attributes drive consumer liking. More importantly, we determined that the fruity and floral attributes of linalool improve the taste of coffee products higher than achievable by varying previously known coffee attributes.

components and traditional drivers. We were surprised by this determination, as one of ordinary skill in the art would not expect a fruity and floral characteristic to enhance the overall liking of traditional, i.e. non-flavored, coffee (see, present specification, paragraphs [0033]-[0040].

- 14. Moreover, an internal Kraft study and corresponding Kraft Research Report, attached in Appendix F, shows that the presence of linalool resulted in a more desirable coffee flavor over the control coffee.
- 15. Referring to the determination of a fruity-floral attribute which drove consumer liking in more detail, the determination was conducted by first identifying groups of naturally occurring coffee chemicals, i.e. coffee components, each of which describe a well-known sensory attribute of coffee.
- 16. In the food art and, in particular, when considering affecting the taste and flavor of a food product, one must be mindful and consider the effect adding one or more flavorant will have on the other compounds or components in the food product to which the flavorant(s) is/are added. As noted above, flavors are often coupled and, in most cases, it is completely unpredictable as to what effect adding one or more flavorant will have on the flavor of the resulting food product. Furthermore, often one does not add just a single flavorant, as typically it will be necessary to add more than one, in order for the resulting food product to have the desired flavor attributes. For example, U.S. Patent No. 4,311,720 (Marmo et al.) discloses a lemon flavored product produced by adding about 21 different ingredients, in order to produce a lemon flavored product.
- One of ordinary skill in the art would not have been led to add linalool to coffee. Linalool, itself, while known to have a fruity-floral attribute, would not specifically

be described as corresponding to any specific flavor, such as lemon (see above, paragraphs 9, 13 and 14). In fact, linalool is more accurately characterized as "sweet floral, petitgrain-like" (see, e.g., <a href="http://en.wikipedia.org/wiki/Linalool">http://en.wikipedia.org/wiki/Linalool</a>, attached in Appendix G). For example, Marmo et al. include it as one of around 21 additional flavor ingredients, in order to produce a lemon flavor. Moreover, one would not describe linalool, itself, as having a lemon flavor. Therefore, should one desire a lemon flavored product, one would not be led to select linalool from among many flavors, which in combination produce a lemon flavor, such as those described in Marmo et al. In order to get a lemon flavor using linalool, one would have to add additional ingredients, as evidenced by Marmo et al.

- 18. More importantly, prior to our discovery, one of ordinary skill in the art would not have known that fruity-floral attributes drove consumer liking of a coffee flavored product. Therefore, one of ordinary skill in the art would not have been led to add a fruity-floral flavor ingredient to a coffee flavored product. It was only after we determined that fruity-floral drove consumer liking, and, in particular, linalool, that one would have any reason to add linalool to a coffee product.
- 19. In addition, even if one would have desired a fruity-floral attribute in coffee, one would not have known that linalool would have produced a desirable coffee product, let alone one which had a "linalool" flavor. Due to the coupling effect of flavor ingredients and the number of different coffee components and flavor ingredients in coffee, one of ordinary skill in the art would not have any reasonable expectation of success or would have been able to predict that linalool would drive consumer liking. Prior to the present invention, it was unpredictable that adding linalool to coffee would be desirable at all.

- 20. Moreover, the claimed amount of linalcol does not produce what one would describe as a lemon flavor or in any way reflect the flavor of linalcol. To the contrary, the coffee product resulting from adding the claimed amount of linalcol would not be described by one of ordinary skill in the art as having a lemon flavor, although it does produce a fruity-floral attribute. Instead, the resulting coffee product would be described or characterized by a consumer as having a natural coffee flavor.
- 21. Furthermore, not only would one not have any reasonably apparent reason to add linalool to coffee, one would not have known how much to add in order to drive consumer liking. Even if one would have thought it desirable to have a fruity-floral attribute to coffee, one would not have been enabled to know how much linalool to add in order to drive consumer liking. For example, if one would have added the 100 parts per 1,000 of flavorant, as disclosed in Marmo et al., one would not have found that the added linalool drove consumer liking at all (see, e.g., present specification, paragraphs [0044]-[0047]. Therefore, even if one would have been led to add linalool, one would not have known to add a sufficient amount of linalool to produce a coffee product with improved consumer liking. Since the amount of linalool of Marmo et al. would not have been considered to improve liking, or change the flavor of the coffee product, one would not be led to increase that amount of linalool to have the claimed amounts. Again, it must be stressed that adding linalool to coffee does not produce a "linalool" flavored product. Therefore, one would not know how much linalool needs to be present in order to improve taste. In other words, the prior art fails to enable one to know to add linalool to coffee, let alone how much to add.

- 22. The amount of linalool added to a coffee product allows one to produce a coffee product which mimics the coffee attributes in high quality, more expensive coffee blends. Accordingly, linalool is added to a coffee product in order to produce a rich flavor which one would find in, or associate with, more expensive coffee blends. However, the linalool does not result in the coffee having what one would describe as a "flavored" coffee flavor. Instead, one would describe the coffee as having a traditional natural coffee flavor (see, e.g., present specification and the Kraft Research Report, Appendix F).
- 23. The undersigned declares further that all statements made herein of his knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of United States Code and that such wilful false statements may jeopardize the validity of the application or any patent issuing thereon.

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Mark Norton

Signed this 20th day of MARCH

# **APPENDIX A**

### A European Sensory and Consumer Study A Case Study on Coffee





The attributes marked \* in Table 5.1 indicate those which were developed by the ICO from extensive previous sensory work on coffee. These were provided to all panels as training aids, and many chose to use the ICO list as a basis for their vocabulary. This aspect clearly affects the occurrence of certain attributes by the panels (see Chapter 2 for details).

Table 5.1 Attributes used by more than one panel, followed by the number of attributes 'unique' to each of the eleven panels, '9 indicates an attribute from the ICO panel.

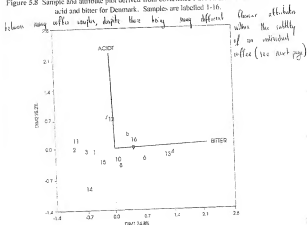
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It was intended to start the construction of a Luropean vocabulary from a fairly simplistic point of view, though to some this may be somewhat controversial. Attributes will be brought together to provide a fuller European sensory vocabulary based on results from this study.

The simple perspective, supported by intuitive reaction to the results of the cluster analysis, was to consider only the attributes bitter taste and acid taste. The rationale behind this was simply that two clusters highlighting both those attributes were apparent: Clusters I and 7 for bitter and Clusters 2 and 4 for acid. Undertaking PCA on these two attributes for the aine panels using both these attributes showed some interesting, though not totally unsexpected, results. Figures 5.8 to 5.16 can be compared with Figures 4.1 to 4.11 to accurate similar fibuse in the sample structures. As can be seen, the overall situatures are very similar fibuse, in general terms the coffee samples can largely be separated on these two basic tastes. However, there is some danger in reading too much into this simplistic view.

Biller (which of reset) and Addity do a good jet of downing the differences.
Figure 5.8 Sample and attribute plot derived from covariance PCA on the two attributes.



# **APPENDIX B**

### Original paper

### Potent odorants of the roasted powder and brew of Arabica coffee \*

forc Blank \*\*, Alina Sen \*\*\*, and Werner Grosch

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intensive Geruchsstoffe von Röstkaffee and Röstkaffeeaufguß aus Arabica-Kaffee

Zewimenfassung, Die Aromaektrakt-Verdünnungsanager (AEVA) von Röstkaffe ergab 13 wichtige Geruchsfeffer: 2-Methyl-3-furanthiol (I), 2-Furfurythiol (II),
Methonal (III), 3-Metapto-3-methylbutylformiat (IV),
2-forcorot-2-methovypvrazin (V), 2-fibryl-3-formiatgerichten (V), 2-fibryl-3-formiatgerichten (V), 3-fibryl-3-formiatgerichten (V), 4-Vinylgunjacol (XII) und
19-framascenon (XIII). Die vergleichende AEVA
12-Rostkaffe und daraus hergestelltem Aufguß zeigte
Abfüguß eine Zunahme von III, IX, Vanillin und 4-Hy2-formiatell (V), VIII und VIII.
2-formiatell (V), VIII und VIII.
2-formiatell (V), VIII und VIII.
2-formiatell (V), VIII und VIII.

immary. Aroma extract dilution analysis (AEDA) redid 13 compounds as important contributors to the
mile of reasted coffee (powder): 2-methyl-3-furanthiol
Eurfurythiol (II), methional (III), 3-mercapto-3rightylyformate. (IV), 3-isopropyl-2-methoxyterior (V), 2-ethyl-3,5-dimethylyprazine (VI), 2-3-diimethylyprazine (VII), 3-isopropyl-2-methoxyimethylyprazine (VIII), 3-isopropyl-2-methoxyimethyl-2(5H)-furanone (XI), 4-vinylguaiscol (XII), and
imasscenone (XIII). A comparative AEDA of the
bowder and brew showed in the brew an increase
DC, vanillin and 4-hydroxy-2,5-dimethyl-3(2H)methyl ad accrease of I, II, IV, V, VIII, and VIII.

Presented in part at the 14th International Conference on Space, San Francisco, USA (July 14-19, 1991)

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nce.to: W. Grosch

#### Introduction

The volatile fraction of roasted coffee has been analysed by many authors (reviews in [1, 2]), who identified more than 700 compounds [3] having a wide variety of functional groups. Attempts to determine those volatiles that actually contribute to the flavour of roasted coffee have been undertaken by Tressl [4] and, recently, by Holscher et al. [5, 6]. On the basis of the odour unit concept [7, 8]. Tressl [4] suggested that 2-furfurvithiol, identified for the first time in coffee by Reichstein and Staudinger [9], was the most important odorant. In addition, he suggested further significant compounds for the flavour of coffee which were also confirmed in the present study. Recently, Holscher et al. [5, 6] using gas chromatography (GC) olfactometry of serial dilutions of the volatile fraction (aroma extract dilution analysis, AEDA [10]), established that some of the odorants suggested by Tressi [4] are indeed intensely involved in the coffee flavour and, in addition, they extended the number of key compounds responsible for the coffee flavour. The aim of the present study was to compare using AEDA the potent odorants of the roasted powder and of a brew prepared from this powder.

### Materials and methods

Coffee. The Arabica coffee (Coffee arabica) from Columbia was supplied by Jacobs Suchard (Remem, FRG.) The coffee beans were medium roasted (3 min) using a Jetzone roaster. The particle size of the roasted and ground coffee was 300–500 µm. The coffee powder was packed in 500 g portions which were sealed under vacuum and stored at -35° C. Coffee powder (20 g) on a coffee-siller was extended with hot water (80–100° C, 500 mil); the brew was immediately cooked in a water-bath at 12° C.

Chemicals. Pure compounds, corresponding to those in Table 2, were obtained commercially nos. 2, 5, 61, 01, 11, 18, 20, 24, 25, 28, 30, and 2-hydroxy-3-methyl-2-cyclopenten-1-one (cyclotene) were from Aldrich (Steinheim, FRG); nos. 1, 7, 23, 32, 38 were from Merck (Darmstadt, FRG); nos. 31 and 34 were from Lancaster (Morceambe, UK); 23-pentandione (no. 3), hexamethyldisilazane

(HMDS) and diphenylterumethyldisilszane (DPTMDS) were from Fluka (Net-Um, FRG); methosal (no. 8) was from Signatude). All the state of the state of

### Syntheses

Bis(2-methyl-3-furyl) disulphide. This was obtained by oxidation of the corresponding thiol according to Evers et al. [12].

Ethyldimethylthiazoles. These were prepared by the Hantzsch condensation of an appropiate  $\alpha$ -halocarbonyl with a reactive thioamide [16, 17]. The MS(EI) spectra agreed with literature data.

5-Ethyl-2.4-dimethyliniazole. The starting material, 3-bromo-2-pentanone was prepared by adding bromine (2.5 g) dropwise at 45-50° C to a mixture of 2-pentanone (8.6 g), water (10 ml) and postsium chlorate (0.15 g). A heavier oil was formed, which was separated from the aqueous layer, washed with 5% sodium blearbonate and water, dried over CaCl, and finally isolated by fractional distillation in a 53% yield.

A mixture of actsmide (0.3 g), phosphorous pentasulphide (0.225 g) and toluene (5 m) in a 59-mi three-needs fund-bottomed flash was heated in a water bath. Where bld trumbed layer formed, 3-brome-2-pentanone (0.4 g) was added droppware to the solvent. After completion of the addition the mixture was refluxed for 1 h, water (1 mi) and cone HCI (0.2 mi) were added and the mixture further heated for 1 h. The solvent was then distilled off through a short-pathway column, the residue was made basic with 50% ageous NoCH and finally the compound was extracted with ether. The or was dried over Na<sub>2</sub>SO<sub>2</sub> and the ether removed by distillation at removed by distillation at removal programs. Practionation yielded (53%) of the target compound.

4-Ethyl-2,5-dimethylthiazole. This was prepared in a 62% yield following the same synthetic route as described above. The starting material for the synthesis of the intermediate 2-bromo-3-pentanone was 3-pentanone.

2-Ethyl-4,5-dimethylthiazole. This was prepared in a 46% yield with phosphorous pentasulphide, propionamide and 3-bromo-2-butanone, which was obtained from 2-butanone.

3-Mercapto-3-methyl-1-butanol. This compound (I, in Fig. 1) was synthesized according to Sweetman et al. [19] using some modifica-

tions. The starting material was ethyl 3,3 dimethylacytic and The sulphur was introduced by reaction with bensythiol 3c the thiochter (IV). The second starting the cleavage of the moiety with Na/NH, (Birch reduction) in order to produce the capto derivative (V), which was then reduced with IAHH, 4782the desired thiolochol (I).

3-Bentylbercapte-3-methylbitumine axis onlysters (IV). A resort bencythin (1.0.5 mo) and city 3-4 dimentylate (0.15 mo) and city 3-4 dimentylate (0.15 mo) and city and city of the city of the solution was cooled and actified with diluted HCI (1.0 mo) axis cooled and actified with diluted HCI (1.0 mo) axis cooled and actified with diluted HCI (1.0 mo) axis cooled and actified with diluted HCI (1.0 mo) axis cooled and actified with distributed (2.5 mo) and the corganic phase was washed with (2.5 mo). And the crysting with MgSO, at 4° C(1 h), the solvening removed by distillation. The residue was then fractionated under pressure through a 25-cm Fisher column to afford pure 3 MC - 11 model MSCCI (1.0 model MSCCI).

3-Mercapite-3-methylbiatanolic acid subjector (V). A 500-m bare necked flask was fitted with a dropping flumel and an insertes in inlet. Ammonia was condensed into the flask at — 78° Canded to the liquid until the solution became blue. The properties were added to the liquid until the solution became blue. The state (V; 100 mmol) was slowly dropped into the solution and the an additional amount of sodium pellets was added. The NTi, are nearowal at soom temperature under a stream of nitrogen durie; 13th period. The excess of sodium was destroyed by addition of the state o

3-Mercapto-3-methyl-1-humon (I). LiAlH<sub>4</sub> (1 mol/L, 50 mL) in a cityl either was placed under an N, atmosphere in a 250-ml three fully either was placed under an N, atmosphere in a 250-ml three fully either (20 ml) was slowly dropped into the mixing which was stirred. The reaction was completed by further stirred with either (20 ml) was slowly dropped into the mixing which was stirred. The reaction was completed by further stirred which was stirred in the reaction was completed by further stirred under refull or 13, was formed. His 500, (10%, with was added to obtain a passon of 1, was formed with interest with dichyl discrete (20 ml) and purplied by column chromatography on siline (24 ml). The 10 ml and purplied by column chromatography on siline (24 ml) (25 ml) (25 ml) (26 ml) (26 ml) (27 ml)

3-Mercapio-3-methylburyljormate (II). The title compound (Fig.1) was prepared by formylation of (1) secording to Stoffelsma et al. [20]. The formylation reagent was prepared by addition of acetabyldride (30 mmol) to formic acid (30 mmol) at 45°C for 2 h in a secrew-caped vial. After cooling with iced water, dry pyriding (3 mmol) was added and then 20 mmol of (1) was slowly dropped.

Fig. 1. Synthetic sequence for 3-mercapto-3-methyl-butarrol (I) and its formic acid ester (II)

This is min into the reaction vial which was cooled with food water. Figuriture was heated for 4 as 44-50° C to complete the formytion and after cooling to room temperature it was diluted with scaley cher (90 m), washed with NaHCO, (34%, wt, 5 x 50 m) and then with brine (2 x 50 m) to neutrality and, finally, dried over \$\frac{1}{2}\text{sol}\$, \$\text{Sol}\$ (0 of 1 is. After enrowal of the solvents the residue was fractionated under reduced pressure through a 25-cm Fisher comun to afford pure material. MS(CD) = 49 (25, M\* + 1), 115 (100, M\* +1-H, S), 103 (40); MS(EI) = 148 (5, M\*), 102 (25), 87 (10), 17 (15), 50 (100), 41 (55).

### Isolation of the volatiles

Examed coffise powder. The roasted coffiee powder (\$0 g) was placed \$7.90 ml two-necked flask and dirthy deterly-protante (2-1), \$\psi\_1\$, \$16 ml) was added. The suspension was gently stirred for 10 min and then frozen for 30 min in liquid N<sub>2</sub>. The flask was adapted to the apparatus described in [21, 22] and the volatiles were distilled off \$0.02 Pa for 3 h. The temperature of the water-bath was then Exercised to 50° C and the distillation continued for a further 2 h. The condensates of the first three cooling traps were combined, fined over Na<sub>2</sub>SO<sub>4</sub> at 4° C and concentrated on a Vigreux column for 3 ml.

Giffe here. The neutral components were extracted from the here 500 mlp with CH, CJ/Fron til 1 (4-1, 1/4, 150 ml) for 15 h using creation perforator (Normag, Hofheim am Taunus, Germany), the extract was concentrated on a Vigreux column (50 × cm) to 100 ml and the volatile fraction was isolated by distillation under Ngh vacuum as described above. The distillate was concentrated by piccydistillation [23] to 2 ml. Thus, the concentration factor was 600 mm (2 ml of 10 ml of

### analytical methods

Series of the management of th

raction D).

\*\*April or formance liquid chromatography (HPLC). For the identification compound no. 14 fraction B was chromatographed with the

column and the apparatus described eastlier [24]. Diethyl ether in pentaun (3+97, v/v) was used as solvent mixture for reparation into five stubfractions that eluted in the ranges 40-65.5 ml (B J), 63-88 ml (B JI), 80-30.5 ml (B III), 10.5-144 ml (B IV) and 14.4-20.0 ml (B V). Compound no. 14 was detected in the subfraction B IV. Eraction D was separated by HPLC using the conditions described in [25] in order to identify compound no. 14.

Capillary gas chromatography (HRGC). This was performed using the glass englishes and fused willide capillaries itsel in Table 1. The glass capillaries were prepared according to Grob [26] using some modifications. The AK (alkal) glass capillary was leached with a solution of 20% HCl (10 h, 130° C), and washed with HCl (1 mol/L, 10 m). The decinication process was followed by a dehydration strey (230° C, 20 min, 2.7 × 10° Pa). The metal-free and dry inner surface was deactivated by persilylation 68, 400° C) using a solution of HMDS/DETMDS/pentane (1+1+2, v/v/v). The capillary was subsequently inside with folusen, nethanot and diethyl ether. The corquently inside with folusen, nethanot and diethyl ether. The cording of the control o

The coated capillary was slowly heated (1° C/min) from 35° C to 250° C (10 h). The quality of the capillary was characterized by the "Grob-Test" [28], the separation efficiency ("Trennzahl") according to Kaiser [29] and the peak symmetry [30]. Some details are summarized in Table 1.

The samples were applied by the "on-column injection" technical, Retention indices (RI) were calculated, and the HRGC conditions were as described in [21]. Precolumns were renounced to avoid adsorption effects by the free silanol groups in the glass connector.

Stability of enoisox compounds during HHGC. An aliquot (a1) of a stock solution of the compound in methanol was injected on each of the right capillaries (Table 1). The peak area obtained was set equal to 100%. The stock solution was diluted with methanol are ported for AEDA [10] and aliquots (same volume as a1) of each divincted solution were analysed by RHGC on the capillaries listed in Table 1. Mean values were calculated of the areas of the peaks obtained for each ocupie or capillaries ("a" and "b") in Table 1, and then projected onto the peak area, which would give the amount dissoved in aliquot a1. These theoretical values were related to the 100% values and plotted versus the amounts of the substance amou

	•			4. 45				
I. Capillaries used for capillary gas tography (HRGC)	HRGC system	Stationary phase, film	Dimension and type of glass	Trennza	hi*	Peak	symmetr	yb
	*,500.00	thickness $(d_i)$	material	C10/11	E10/11	ol	ď	Cu
	Ia	SE-54 d <sub>1</sub> =0.3 μm	25 m×0.3 mm Soft glass	40	32	1.1	1.5	1.0
	ľb	SE-54 4 = 0.3 μm	30 m × 0.3 mm Soft glass	40	30	1.1	1.3	1.0
inzahl (separation efficiency) accord-	IIa	OV-1701* d=0.25 mm	60 m × 0.3 mm Fused silica	55	40	1.0	1.0	1.0
Calser [29] using the GC conditions of [26]. C <sub>10/11</sub> , decane/undecane; methylester of decanoic acid and	ПР	OV-1701 d <sub>1</sub> =0.3 μm	30 m × 0.3 mm Soft glass	44	35	1.0	1.0	1.0
tion acid  1 year netry according to Meyer [30].	Ша	Carbowax <sup>d</sup> , $d_t = 0.25  \mu m$	30 m × 0.3 mm Fused silica	27	32	1.1	1.1	1.1
manol: D, 2,3-butandiol; C11,	шь	Carbowax d d <sub>c</sub> =0.25 µm	30 m × 0.3 mm Fused silica	28	-31	1.0	1.0	1.0
(J & W) obtained from (Hofheim, FRG)	<b>IVa</b>	FFAP** d=0.25 um	30 m × 0.3 mm Fused silica	30	,30	1.0	1.0	1.2
Supelcowax 10 obtained from Til Sulzbach, FRG)	IVb	FFAP*** 4=0.25 μm	30 m × 0.3 mm Fused silica	25	31	1.0	1.0	1.1
and buase								

### Mass spectrometry (MS)

MS analyses were performed on au MS 8230 (Finnigan MAT). Bremen, FRQ) in tandem with the gas chromatography (GC) capillary columns described above using the same FRGC conditions. MS(EI) were generated at 70 eV and MS(CI) at 110 eV with isobutane as the reactant gas.

Gas chromatography/olfactometry (HRGC/O)

Aliquots of the volatile fractions were separated by HRGG tailed in Table 2 and odorants were perceived at a smilling por 21]. The sensory significance of each odorant was evaluated apressed as the flavour dilution (FD) factor [10, 24], "thresholds in air were determined by HRGG/D [10, 21]."

Table 2. Potent odorants (FD factor ≥ 16) of the roasted powder and brew of Arabica coffee

No.	Compound	Frac-	rac- Retention index on			Aroma quality <sup>b</sup>	FD fac	tor*	Sensory significa establis earlier	
			OV-1701	SE-54	FFAP		Powder Brew			
1	2,3-Butandione* (diacetyl)	В	686	580	990	Buttery	16	32	[S] [4], [S] [4], [5]	
2	3-Methylbutanal*	В	739	650	950	Malty	16	32	[4]. [5]	
3	2,3-Pentandione	В	791	695	1060	Buttery	32	32	[4], [5]	
4	3-Methyl-2-buten-1-thiol	A/B	874	821		Amine-like	32	< 16	121	
5	2-Methyl-3-furanthiol	A	930	870		Meaty, boiled	128	< 16	[5]	
6	2-Furfurylthiol*	Α	1004	913	1440	Roasty (coffee-like)	256	64	[4], [5]	
7	2-/3-Methylbutanoic acid*	D	1022	860		Sweaty	64	64	[5] [5]	
8	Methional	c	1040	909	1455	Boiled potato-like	128	512	[5]	
9	Unknown	D	1073		1365	Fruity	32	16	-	
10	Trimethylthiazole*	В	1074	997	1370	Roasty, earthy	16	< 16	-	
11 12	Trimethylpyrazine*	D	1078	1000	1395	Roasty, earthy	64	32	[5]	
13	Unknown	C D	1107	1055		Roasty, sulphury	128	. 32	-	
13	3-Mercapto-3-methyl-1-butanol*		1127	972	1655	Meaty (broth)	32	64	[5]	
15	3-Mercapto-3-methylbutylformate*	В	1138	1023	1517	Catty, roasty	2048	256	[5]	
16	3-Isopropyl-2-methoxypyrazine* 5-Ethyl-2,4-dimethylthiazole*	В	1146	1097	1428	Earthy, roasty	128	32	[5]	
17		D	1149	1078	1435	Earthy, roasty	32	16	-	
18	2-Ethyl-3,5-dimethylpyrazine* Phenylacetaldehyde*	D	1154	1083	1453	Earthy, roasty	2048	1024	[4], [5]	
19	Unknown	B/C	1178	1053	1635	Honey-like	64	32	. [5]	
	Linalpol*	C/D	1185	1103		Roasty-earthy	128	128	-	
21		ç	1193	1102		Flowery	32	< 16	[5]	
22	2,3-Diethyl-5-methylpyrazine	C	1218	1155	1485	Earthy, roasty	512	128	-	
	2-Hydroxy-3,4-dimethyl-2-cyclo- penten-1-one	_	1226	1075	1840	Caramel-like	64	128	-	
23	Guaiacol*	Ç,	1228	1093	1850	Phenolic, burnt	32	16	[4], [5]	
	4-Hydroxy-2,5-dimethyl-3(2H)- furanone* (HDF)	D '	1235	1065	2035	Caramel-like	16	256	[4], [5]	
25	3-Isobutyl-2-methoxypyrazine*	B/C	1237	1186	1520	Earthy	512	128	[4], [5]	
26	Unknown	C	1254	1184		Roasty, earthy	512	32	1-35 1-3	
27	5-Methyl-5(H)-cyclopenta[b]pyrazine*	В	1260	1145		Roasty, sweet	32	< 16	[5]	
28	(E)-2-Nonenal*	В	1275	1160		Fatty		< 16	[4], [5]	
29	Unknown	D	1305		2090	Caramel-like	16	16		
30	3-Hydroxy-4,5-dimethyl-2(5H)- furanone* (Sotolon)	D	1347	1107	2200	Seasoning-like	512	2048	-	
	4-Ethylguaiacol*	С	1424	1287	2032	Spicy	256	512	[5]	
	p-Anisaldehyde*	B/C	1431	1263	2030	Sweet, minty		< 16	-1-1	
33	5-Ethyl-3-hydroxy-4-methyl-2(5H)- furanone*	D	1433	1193	2270	Seasoning-like	512	1024	=	
4	4-Vinyiguaiacol*	С	1482	1323	2205	Spicy	512	512	[4], [5]	
55	(E)-β-Damascenone*	В	1502	1395	1815	Honey-like, fruity	2048	64	[5]	
6	Unknown	B/C	1620		2355	Amine-like	64	64	[2]	
57	Bis(2-methyl-3-furyl)disulphidef	A	1640	1540	. 2150	Meaty, sweet	32	128	-	
8	Vanillin*	D	1645	1410	> 2500	Vanilla-like	32	512	-	

Fraction in which most of the compound appeared after enrichment by column chromatography

Odour description assigned during aroma extract dilution analysis (AEDA)

<sup>\*</sup> The flavour dilution (FD) factor of the compounds was evaluated using the capillaries given in brackets: no. 5 (SE-54), nos. 7, 13, 22-24, 30, 33, 38 (FFAP), the resting odorants (OV-1701) and the sensory significance of the compound for the flavour forested coffice (powder) was reported by the quoted authors

<sup>•</sup> The compound was identified by comparing it with the reference substance on the basis of capillary gas chromatography (HRGC) on the capillaries presented in the table, the mass spectrum and the odour quality and threshold, which was perceived at the sniftling.

The mass spectrometry signals of the substance were too weak for an interpretation; the compound was identified by comparing I with the reference substance on the basis of the resting criteria reported in footnote.

### Results and discussion

### Stability of enoloxo compounds during HRGC

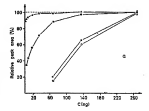
Sifice contains 4-hydroxy-2,5-dimethyl-3(ZH)-furanone (TDF) and other compounds having an enolox sub-furcture [4, 5, 14]. HDF is partially degraded during malysis by HRGC [31], which would affect the results of AEDA. Therefore, in preliminary experiments, the yields of HDF and cyclotene, as examples for enoloxo compounds, after HRGC in relation to the amounts injected and GC capillaries coated with four stationary phases (Table 1) were determined. To calculate the yields of the dilited samples, the peak areas were projected on to that full the control of the stationary phases (Table 1) were determined. To calculate the yields of the dilited sample, which was set 100%. Figure 2 indicated that during HRGC on the relatively unpolar capillaries SE-54 and OV-1701 the yields of HDF and Spelotene decreased strongly with increasing dilution of the sample. HRGC on Carbowax improved the results but the highest yields were obtained on the capillary Gated with FFAP. Consequently, this capillary was used for the AEDA of the enoloxo compounds.

### Odorants of the roasted powder

AEDA of the coffee powder revealed 38 odorants with ED factors of 16 or higher. These compounds were entitled by column chromatography on silica gel as deficiled in Table 2. Compounds nos. 14 and 17 were entitled further by HPLC of column fractions B and D, respectively. HROC-MS analysis and HROC/O fractions A-D and the subfractions obtained from HPLC resulted in the identification of 28 odorants (Table 2). Four simpounds (nos. 4, 5, 21 and 37) gave unclear MS significant processes of the criteria respect of the control of the column of the c

In the higher FD factor range 128 to 2048 (Table 2) to Dolowing 13 key compounds of coffee flavour were controlled to the controlled to t

J. AEDA the authors [5, 6] diluted the extract 500mind nine odorants at this level and identified eight in. The potent odorants nos. 5, 6, 8, 14, 25, and 35 mong them as well as 2-/3-methylbutyric acid (no. HDF (no. 24), which showed relatively low FD and our study (Table 2). Holscher et al. [5, 6] resorting the RI data (219) and 2257 on DB-Wax) retion of the RI data (219) and 2257 on DB-Wax) re-



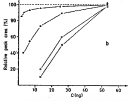


Fig. 2a, b. Yields of 4-hydroxy-2,5-dimethyl-3(2H)-furanone (a) and cyclotene (b) after the capillary gas chromatography (HRGC) on capillaries SE-54 (o), OV-1701 (n), carbowax (•) and FFAP (Δ); C, compound; cf. Table 1

ported [5, 6] we suggest that these odorants were identical with sotolon (no. 30) and EHMF (no. 33), respectively. Sotolon was also been found in sherry [32]; cane sugar [33], fenugreek seeds [34], and aged sake [35], while EHMF has been detected in acid-hydrolysed vegetable proteins [36, 37].

The conclusion of Tress [4] that compounds nos. 2, 3, 6, 17, 23, 24, 25, 28, and 34 play an important role in coffee flavour was supported by AEDA, in particular for nos. 6, 17, 25, and 34, which showed higher FD factors (Table 2). Kahweofuran, which was also proposed as an impact compound of coffee flavour [4], was indeed identified (data not shown), but its very low FD factor suggested that it did not contribute to the coffee flavour.

### Odorants of the brew

A brew was prepared from the powder of the Arabica coffee sample. Its odorants were extracted and evaluated by AEDA. As the FD factor of a compound is proportional to its concentration in the extract [10] the results listed in Table 2 suggested that, compared to the powder, methional (no. 8), HDF (no. 24), sotolon (no. 30) and

vanilin (no. 38) increased in the brew. On the other hand the thiols nos. 5, 6, 14 and the pyrazines nos. 15, 21, 25 decreased strongly. These changes in the concentrations of the odorants might be caused by a lower solubility in the brew, as was found for (E)-B-damascenone [22], and by degradation of the odorants by the hot water used for preparation of the brew. In addition, the FD factors of water-soluble odorants (e.g. diacety), 2,3-pentandione) might be reduced by a low extraction yield from the brew.

### Odour thresholds

The odour thresholds in the air of some key compounds of the coffee flavour were evaluated (Table 3). The lowest threshold was found for 3-mercapto-3-methylburylformate. It was 50-fold lower than the threshold of 2-fur-furylthiol, the impact compound of the roasty odour note of the coffee powder. The threshold of 2-ethyl-3,5-dimethylpyrazine was in the same range as that of the 2-fur-furylthiol. Together with (B)-f-damascenone, the threshold of which was four-fold lower, this pyrazine appeared with the highest FD factor in the extract of the coffee powder and was also important for the flavour of the brew, the door thresholds increased in the order 5-ethyl-3-hy-droxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-5-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-methyl-2(SH)-furnaone, stotlon, 2-hydroxy-4-meth

Table 3. Odour threshold of some volatiles identified in coffee (powder and brew)

Compound	Threshold (ng/L; air)	Capil- lary
3-Mercapto-3-methylbutylformate	0.0002-0.0004	OV-1701
3-Isopropyl-2-methoxypyrazine	0.0005-0.001	OV-1701
2-Methyl-3-furanthiol	0.001 -0.002	SE-54
3-Isobutyl-2-methoxypyrazine	0.002 -0.004	OV-1701
5-Ethyl-3-hydroxy-4-methyl-2(5H)- furanone	0.002 -0.004	OV-1701
(E)-B-Damascenone	0.002 -0.004	OV-1701
2-Ethyl-3,5-dimethylpyrazine	0.007 -0.014b	OV-1701
2,3-Diethyl-5-methylpyrazine	0.009 -0.018*	OV-1701
2-Furfurylthiol	0.01 -0.02	OV-1701
Sotolon	0.01 -0.02	FFAP
4-Ethylguajacol	0.01 -0.03	OV-1701
2-Hydroxy-3,4-dimethyl-2-cyclo- penten-1-one	0.05 -0.1	FFAP
Methional	0.08 -0.16	OV-1701
4-Vinylguaiacol	0.4 -0.8	OV-1701
Trimethylthiazole	0.5 -1.0	OV-1701
Vanillin	0.6 -1.2	OV-1701
4-Hydroxy-2,5-dimethyl-3(2H)- furanone	0.5 -1.5	FFAP
3-Methylbutanal	2 -4	OV-1701
Diacetyl	1020	OV-1701
2,3-Pentadione	10 -20	OV-1701

Capillary used for the determination of the odour thresholds by HRGC/olfactometry

3,4-dimethyl-2-cyclopenten-1-one and HDF. Dimerand 2,3-pentandione were important odorants showled relatively high thresholds.

#### Conclusions

The character impact odour compounds of the postand brew of Arabica coffee are different. The contritions of thiol odorants (e.g. 3-mercapto-3-methylbhi formate, 2-furfurylthiol) and (E)-β-damascenons stronger in the flavour of the powder than to that of brew. The reverse effect was found for methional, allon, 4-hydroxy-2,5-dimethyl-3(2H)-furanone and valin. 2-Ethyl-3,5-dimethylpyrazine plays an important in the flavours of both the powder and the brew.

Acknowledgements. This work was supported by the Forschung kreis der Ernährungsindustrie (Bonn) and the AIF (Köln). Wegrateful to Miss Foschum, Miss Kustermann, and Miss Relaham for skilfull technical assistance.

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b The thresholds were determined by C. Cerny (private communi-

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# **APPENDIX C**

WILEY

FLAVOR CHUNDERR

**^**сно

Ivon Flament

### **COFFEE FLAVOR CHEMISTRY**

### Ivon Flament

formerly of Firmenich SA, Geneva, Switzerlan

With the collaboration of Yvonne Thomas-Bessière

For the best part of two centuries investigators have tried with varying degrees of success to leanlify the compounds which give reasted coffee its characteristic arona and taste. The analytical methods and the state of progress in chemistry at the end of the 19th century did not allow for the separation, isolation and felentification of the multitude of trace chemicals which are present in reasted coffee. By 1900, scarcely a dozen compounds had been identified. Since the beginning of the sixties, with the advent of gas chromatography and mass spectrometry, the rumber of identifications has increased tremendously. To date, 850 compounds have been identified in the flavor of roasted coffee and 300 in the small of green coffee.

In this work, the authors systematically review the non-volatile constituents of green coffee, including their structure, and discuss their important contribution as flavor precursors during the roasting process. They also trace the chronological discovery of the individual chemicals and critically examine the validity of their identification, highlighting the enormous progress which has been realized during the twentieth century and particularly in the tast 40 years. For convenience, the constituents of green and roasted coffee have been distributed into chemical classes according to structure, systematic and empirical names, their CAS Registry Numbers and occasionally their FEMA classification. Comments are made on the origin or the formation during roasting of each Individual compound.

### Coffee Flavor Chemistry:

- contains an up-to-date list of almost 1400 original literature references;
- is the first book to provide a comprehensive overview of coffee flavor chemistry;
- critically discusses all of the identified and confirmed compounds in coffee;
   presents the major part of the book as a catalogue, for the benefit of the reader;
- Includes Information on structures, systematic and empirical names, identification, mechanism of formation, synthesis, detection threshold and organoleptic properties of each constituent where available;
- devotes a chapter to the flavor precursors, including the names and structures of the compounds with reference to the corresponding analytical work.

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This alcohol has been identified as a degradation product of sotolone (G.12) in the presence of UV light (Martin  $et al_n$ , 1990).

The flavor is fusel, solvent, estery, leafy (Chemisis, 1999)

(B.31) (B.31) 1-Octen-3-ol. oct-1-en-3-ol. amyl vinvl carbinol, pentyl vinyl carbinol, 3-hydroxy-1-octene. Matsutake alcohol [3391-86-4] FEMA 2805; (±)- [50999-79-6]; (R)-(-)- [3687-48-7]; (S)-(+)- [24587-53-9]

Identified by Stoll et al. (1967) in roasted coffee, by Cros et al. (1980) (headspace analysis) and in green coffee by Gutmann et al. (1979). In green coffee, the concentration is 0.03 ppm for Holscher and Steinhart (1992) and the peak area in GC represents 0.60% of the volatiles for Cantergiani et al. (2001) (vaccuum hydrodistillation).

It is one of the most intense flavor compounds formed by autoxidation of linoleic acid  $(C_{18,2})$  (UBrich and Grosch, 1987). As an aside, it is interesting to note that in mushrooms, an enzymic oxidative breaktlown of linoleic acid gives the (R) isomer by the intermediate of 10(S)-hydroperoxy-8(E), 12(X)-octadienoic acid (Grosch and Wurzenberger, 1985).

These authors mention a mushroom aroma and give an odor threshold range of 2.3.5.3 ppb determined by high-resolution GC olfactometry. The (S)-isomer is described with a green, vegetable mouldy flavor (Chemiss, 1993) and the (R)-isomer with a green, unshroom meart (flavor (Chemiss, 1992).

(B.32) 7-Octen-4-ol, oct-7-en-4-ol. 1-octen-5-ol [53907-72-5]; (±)- [87830-31-7]

Identified in green coffee by Gutmann et al. (1979) and found by Guyot et al. (1982, 1983) in healthy us well as 'stinking' green coffee.

It is characterized by a powerful, earthy note.

(B.33) 5-Hepten-2-ol, 6-methyl-, 6-methylhept-5-vn-2-ol [1569-60-4]; (±)- [4630-06-2]; (R)- [58917-27-4]; (S)- [58917-26-3]

Identified by Cantergiani et al. (2001) in a green Mexican coffee (0.25% of the volatiles by GC). The corresponding ketone (D.33) has been identified but only in roasted coffee.

The racemic mixture has a green, fatty odor (Chemisis, 1981).

(B.34) (B.34) 3-Buten-2-ol, 2-methyl-, 2-methylbut-3-en-3-ol, 1.1-dimethyl-2-propenol, 1.1 dimethylallyl alcohol, dimethyl vinyl carbinol, 3-hydroxy-3-methyl-1-butene [115-18-4]

Identified by Silwar (1982). Silwar et al. (1987) give a concentration of 0,20-0,35 ppm in roasted coffee steam distillation and distillation and most only on the confee steam distillation and most only in a reasted antiblea. disappearing upon producing decadancy and not in the various green arabicas and robustes that they examined. Cantergrain et al. 2001) identified it in a green Mexican coffee where it represents 0.13% of the valatiles by GC fafter vaccuum hydrodicillation).

The odor and flavor are described as sob ent (Chemisis, 1972).

Lindock

(B.85) (B.85) [1.6-Ociadien-3-ol, 3,7-dimethyl) 3.7-dimethylocus/1.6-dien-3-ol, 2.6-dimethyl-2.7-ociadien-6-ol, hindod [B-70-6] [EBAA 2635] (b) [22564-99-d]; ficureol. (B)-(-) [126-91-0]; coriandrol. (S)-(+) [126-91-9]

Identified by Stoll et al. (1967). Stoffelsma et al. (1968), and Friedel et al. (1971) (IR. MS data given) in roasted coffee. Ho et al. (1993) gave a concentration of 0.73 ppm in a roasted Columbian coffee (head-

indayable

ul ingelies

spacer. Linabool was found by Guyot et al. (1982, 1983) in 'striking' green coffee, but not in a bealthy variety. On the contrary Spadone et al. (1990) found at in green beams of a Puerro Rico 'Rio' and of a healthy variety. Helscher and Steinhart (1995) give a concentration of 0.175 ppm in green coffee distillation extraction at room temperature then at 70 C. It represents 1.16 (2007 of the volatiles scramme hydrodistillation at room temperature in the analysis of a green Markian arabica by Canterglant et al. (2004). Linabool has also been identified in red berries of an arabica (grown in a greenfulness), the amount decreasing with time with a concomitant mercase of finabool voides (see 1.56, 1.57, 1.144 and 1.145).

Landsof has a light and reiseding, than layout, edge with a faintly cities; not extratable, 156-11, the or of the racements would, fortal press beganning them as 1991. The typical borated means of the properties of a rabbia coeffect on not of the properties of the

### Diols and ethers

(B.36) Ethane, methoxy-, methoxyethane, ethyl methyl ether [540-67-0] Identified in headspace analysis of roasted coffee by Wang et al. (1983).

(B.37) Ethane, I.I. oxybis-, ethoxyethane, diethyl ether, ethylether, diethyl oxide [60-29-7] Identified in four green arabicus (out of six) and in five green robustus (out of six) by Procida et al. (1997) who did not find it in a rousted arabicu (headsmace, GC/MS).

(B.38) Propane, Z-ethoxys. 2-ethoxypropane, ethyl isapropyl ether [625-54-7] identified by Ho et al. (1993) in a headspace of reasted Columbian coffee, with a concentration of \$3.8 ppm

(B.39) Propane, 2-methoxy-2-methyl-, 2-methyl-, 2-methylpropane, vert-buryl methyl ether, 1.1-dimensylethyl methyl ether [1634-04-4]

identified in headspace analysis of roasted coffee by Wang et al. (1983).

Crayot et al. (1983) mention the presence of dimethoxycyclobexane, without any structural precision , in stinking as well as in healthy green beans.

(B.4b) 2.3-Butanediol, Intana-2.3-diol, dimethylethyleneglycol, 2.3-binyleneglycol [\$13-85-9]; (R\*,R\*) [\$5007-63-7]; (2R.3R) (or [R.4R\*,R\*)]+10 r [k.0c, or three, 2R.3R) [24347-58-8]; (25.35)+(or [S-(R\*,R\*)]+1) or divite, or three, 2S.3S) [1913-06-0]; (R\*,S\*)+(or mess, or erythree, 2R.3S) [\$341-95-7] Identified by Vincent et al. (1976) in sinding arosa bears, the great proceedings of the processing and processing arosa bears. It is proceeding the processing arosa bears. It is processed to be proceeding the processing arosa bears. It is processed to be proceeding the processing arosa bears. It is processed to be proceeded to be pr

Identified by Vincent et al. (1976) in stinking green beans. It was noted as butane diol-2, in the text and in the semmary, but with the presence of the corresponding diference and hydroxyletone in the same

Methyl propanoate has a very diffusive aftereal-run-like odor. Sweet and fruity of very poor tenacity (Arctander, 1967). The flavor of fruity, greefi, chemical (Chemica, 1997). Ahmed et al. (1978) gave a probable door threshold in water of 58 ppb.

### (F.26) Propanoic acid, propyl ester, propyl propionate, propyl propanoate [106-36-5] FEMA 2958

Identified in a green coffee headspace analysis by Merritt et al. (1970), only in a Columbian arabica and not in a Statios trabica and a robusta.

Firsh-athereal, fruity-floring does of very poor tenacity. The fruity notes being apple-pincapple-like, with other branch haracteristic floring of a general fill of sweetness (Arctander, 1967). The flavor is othereal, fruity, Spin, weak (Chemiss, 1998). The odor threshold given by Father or J. (1967) was \$75 per.

(F.27) 2-Propatione, 1-(1-oxoprupoxy). 2-axoprupyl proplantae. 2-axoprupyl proplantaele, hydroxy accione propionate, acetol propianate, acetonyl propataente [72845-79-5]. Identified by Bondarovich et al. (1967) in an 'aroma complex' of roasted coffee.

### (F.28) Butanoic acid, methyl ester, methyl butyrate, methyl butanoate [623-42-7] FEMA 2693

Identified in roasted coffee but absent from green beans according to Mercitt et al. (1970).

This ever has a serie diffusive and penetrating, sweet-ethereat, fruity odol. In extreme dilution it is requisited in a sipple peel both a slightly fatty peach-like undertions (Aroundler, 1967). It has family over-type, theory that of Continued (Continued of the Continued Continued of the Conti

### (F.29) Butanoic acid, ethyl ester, ethyl butyrate, ethyl butanosate [105-54-4] FEMA 2427

identified by Merritt et al. (1970) in a green coffee headspace of a Columbian archica but not in a. Santos archica and a robustu coffee. It has also been found by Guy of et al. (1982, 1983) in stinking green coffee, and in a very low concentration in healths beans of a Courseron archica.

Its pine-cone character can exemple and very diffusive off-flavor. Powerful, ethereal-fruity oldor suggestive of busing land pineapple and very diffusive (Arctander, 1967). The theory is fruity tresh (Chemiss, Surface).

An odor threshold of Lppb is given by Flath et al. (1967). The flavor threshold was 0.45 ppm in water too Kesh and Powers (1968) and 15 ppb for Sick et al. (1969). Ahmed et al. (1978) gave lower vidues modelife edor threshold inwater 0.15 ppb floor with confidence firms (0.003–4.6) and flavor threshold et al. (1978) gave flower vidues to 1975 ppb with confidence firms (0.0008–2).

### flutumeic seid, otheryl ester, vinel bisamouso [123-20-6]

This was sentalizely identified in headspace of row or cornelly. Wang confidences

### el "In». Pentannic acid, methyl ester, methyl je unassace, methyl valerate [624-24-8] FEMA 2752

H- provides at headspace of green coffee, but not or toasted beans, was mentioned by Merriti et al., 1970.

is it is discribed as having a pungum ethoreal Green-trials applicable actor of poor terricins (An trader. 1907).

Of ~ 1150 compromeds within this reflection feet year many house failing or floral -type characters associated with them

The hidici hall tous linears

(F.M) Pennanic acid, othyl ester, ethyl pennanoute, ethyl valerace, ethyl valeriaidde [539-82-2] FEM 4-2462

Identified by Merritt of all (1970) in the headstrace of green beans of a Collambian architecture in a Sames and the a to-buston.

The solid processed and diffused others bright materials will a term assessment a frequency frequency from the first of th

(F.32) Pentanoic acid, buryl ester, but el protonoute (butyl suicrate, buryl videriona); [501-68-4] Fh.M 4-22 F

Idemard in given becaus obtailes simulatineous distillation extraction and Groot the concentrated by Spindom and Lindom 1997 could in Paerro Bloot Rev coffee and not in other Rich and freshing beaus. The Broot is frestly, appreciation of themass, 1978).

(F.33) (F.33) Hexanole acid, methyl ester, methyl hexanoute, methyl caproate [106-70-7] FEAIA 3708

Identified by Merrits of al. (1970) in green and roasted beans.

It is described as having a powerful ethereal and diffusive, weet odor of piscapple (prico) type (Aretander, 1963).

(F.34) Hexanoic acid, ethyl ester, ethyl hexanoute, ethyl capcoure [123-66-0] FEMA 2439

Identified in green immarate beans by Full et al. (2000), after electronic sorting of the beans followed by simultaneous distillation extraction—

It is described as procappie, banana (fruity) The odor threshold in water was 1 pph according to Takeoka et al. (1995).

(F.35) Terradecanoic acid, 4-methylethyl ester, isopropyl retradecanoute, isopropyl myristate [110-27-0] FEMA 3556

Identified by Ramos et al. (1998) in headspace after solid-phase unicroextraction of a browed analyca (compare with F.8 and F.19).

(F.36) (F.36) Hexadecanoic acid, methyl ester, methyl palminate, methyl hexadecanoate [117-39-0]

Identified by Stoff et al. (1967) in a roasted coffee extract and by Cantergiani et al. (2001) on a green Mexican arabies extract after vacuum hydrodistillation) where it represents 0.04% (GC)

(F.3") Hexadecanoic acid, ethyl ester, ethyl publicate, ethyl hexadecanousc [698-97-7]

(F.38) Hexadecanole acid, butyl ester, hayl palmirate, butyl besadecanoste [111-06-8]

Those have been found, like the lower homolog (see F.36), by Cantergiant code, (2011) in a green Mexican codies (respectively 0.12 and 0.05% of the extract in GCi. F.37 had previously been tenuto ely identified by Spadone and Liardon (1988) in 'Rio' and healthy green beans.

The flavor of F.38 is described as facty, waxy, mouthfeel (Chemisis, 1989).

# **APPENDIX D**

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stiffing perc, abber trained people evaciates and teacor and a heasts stiffing perc, abber trained people evaciates and record the essency inversion of individual components (Hodderter et al., 1990). Distant rechniques are used to determine the se-celled finerior (HD) factors. By a suspires distarion of the amount secure (111 by volume) followed by CCO analysis the most important countibutors can be srateled at the highest distant, thus getting the highest ED factors. Dist extending the developed by Ultical and Groods (1985), is known as arounversed as the concentration and Groods (1985), is known as arounversed as the concentration whiched by its factorial deals of represent a few concentration whiched by its factorial deals of strength and the colour introduled of a subsance are known. A method for the prediction of OAV from BD factor is reported as well as the precision and the vertical design of KBDA (Ferreira et al., 2001).

# 4.4.4 Aroma impact compounds in roasted coffee

In arabica coffee the most important contributors to the around of roast and ground coffice are determined by the techniques mentioned above.
3-Mercupor—benchyllenyforment (MMISP) Zunfurjhlioti, unchfamil, and Ammacenone and the two preparation and funniones show the highest FD factors (Hodecher et al., 1992). A comparison of the study of the proper exactly good specification is given in Table 4.5, together with the around impressions of the substance.

The abother amounts of the arona compounds shown in the last command of the count of the arona compounds shown in the last of the alphur compounds, is demonstrated by their law concentrations of the sulphur compounds, is demonstrated by their law concentrations are agreemed with a compound of the compounds of the alphur are afficient to generate the account support of the aronagest colour impression, which is rone at the aronagest colour impression, which is rone of 17000.

# 4.4.5 Effects on cup impression

Everybods who has ever laid a really had our of coffee in direct experience on perfect on Renow being the difference can be. The perceived quality depends on objective criteria such as quality of green force, but also on subjective or cultural preferences like type of preparation.

4-Vinylguaiacol

1280 952 5Z 500 Clove-like 4-Ethylguaiacol 3450 \$201-Z15 Phenolic, burnt loseleud 852 821-91 1054 005 Honey-like, fruity enonessemed-q-(3) 000 SZ 1054 19 009 Yreaws 3-Mercapto-3-methylbutanol 2/3-Methylbutanoic acid 100 52 20 20 20 20 zε any-dnog 0052 HOUGH-like **p**9 -18 Peanut-like V9-V ZIS Paprika-like 051 1074 800Z 005 007 5 007 005 Catty, roasted coffer ες Yosely, musty 001 Vate 1 (E)-2-Nonenal SS 1054 8102 Visum , Vissoon 2-Ethyl-3,5-dimethyl-pyrazine 851 1024 Cooked potato 821 Isnonal Vinegar-like Acetic sold 158 100 200 200 200 200 500 **Asea**d S-Isopropyl-3-methoxy-pyrazine 1320 1054 526 toasty, coffee-like lointhruthut-S 0009 7F-01 Visum , Viseo F **b9** 2,3,5-Trimethylpyrazine Sweety, cetty Mercaptopentanone anoibanetnaq-E,S. irl}-1-natud-S-lythaM-E ioirtinesul-E-lythaM-S anonetoendrasses n9 35 158 Roasted mear-like 13 957-79 -Animal-like, skunky 001 00585 871-Þ Buttery 00455 7201-95Z gnifeir 2-Methylpropanal 2-Methylbutanal 2,3-Butanedione 28 600 24 000 87L-91 Pungent, fermented 001 52 Pungent, malty 0050 Putrid, cabbage-like Methanethiol wayer\* (µg/kg) Schenker<sup>3</sup> (FD) Grosch<sup>2</sup> (FD) Holscher<sup>1</sup> (FD) Odour impression Increar

Sources: "Holscher et al. (1990); "Czerry and Grosch (2000); "Schenker et al. (2002), range according to roassing isothermal high/low temperature;

500

35

215

Table 4.5 Aroma impact compounds in roasted coffee powder

Peaks sorted by retention time on D8-wax column. Compounds with FD ≥ 32 are reported.

Vanilla-like

# **APPENDIX E**

### Analysis of Roasted Coffee Powders and Brews by Gas Chromatography-Olfactometry of Headspace Samples

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The highly volatile, potent odorants of coffee samples were evaluated by gas chromatography-olfactometry of decreasing headspace samples (GCO-H). 2,3-Butanedione, 2,3-pentanedione, 3-methyl-2-butenthiol (I), methional, 2-furfurylthiol (II) and 3-mercapto-3-methylbutylformate (III) were the key odorants of both, the powders of Arabica and Robusta coffees. 2-Methyl-3-furanthiol (IV), 2,3-diethyl-5-methylpyrazine and an unknown compound were additional key odorants of the latter. An increase in the odour potencies of acetaldehyde, propanal, methylpropanal, 3-methylbutanal and dimethyltrisulphide as well as a decrease in the odour potencies of the thiols I to IV in the brews were the major differences with regard to the powders.

### Introduction

Odorants contributing to the smell of freshly roasted coffee beans have been screened by aroma extract dilution analysis (AEDA) and then identified (1.2). However, AEDA is limited to odorants boiling higher than the solvent used for the extraction and dilution steps. Furthermore, odorants boiling in the same range as the extraction solvent are partially lost during the concentration of the extract by distilling off the solvent. To overcome these limitations, AEDA was completed by gas chromatography-olfactometry of headspace samples (GCO-H) (3,4).

The highly volatile odorants of freshly roasted coffee beans have been screened by GCO-H (3), and, by analogy with AEDA (5) the results have been expressed as flavour dilution (FD) factors. The FDfactor for a compound is a measure for its odour potency and is defined as ratio of its concentration in the largest headspace sample to its concentration in the smallest one in which odour was detected by GCO (3). In the headspace of coffee 12 odorants were identified in the largest sample of 250 µL (3). Methanethiol and 2-methylpropanal showed the highest FD-factor of 25; 2,3-butanedione, 2,3-pentanedione, 3-methylbutanal and 2-furfurylthiol followed with an FD-factor of 5. Headspace samples often contain odorants in such low concentration levels that they give no detector response. In the case of coffee, 2-furfurvithiol, 3-methyl-2-butenthiol and methional are examples which are only detectable by sniffing of the GC effluent (3). However, as discussed recently (4), most of these odorants can be identified in headspace samples on the basis of the results obtained by a preceding AEDA. In the present study, GCO-H was applied to compare the highly volatile odorants of the powders and brews prepared from roasted Arabica and Robusta coffees. In addition, a brew obtained from a soluble coffee powder was included in the comparison.

### Materials and Methods

The Arabica coffee (Coffea arabica) from Columbia and the Robusta coffee (Coffea canephora var. robusta) from Indonesia were medium roasted and then ground (particle size: 300-500 µm). The coffee powders were packed in 1 kg portions which were sealed under vacuum and stored at -35°C until use. Hot water (1.1 L. ca. 95 °C) was poured on the coffee powder (54 g) in a filter (Kaffee-Filterpapier Nr. 4, Plus Warenhandelsgesellschaft, Hamm, Germany) yielding 1 L of the coffee brew. A soluble coffee powder was purchased from a local market; it (16 g) was dissolved in 1 L of hot water (ca. 95 °C).

Pure compounds, corresponding to those in Table 1 were obtained commercially from the sources (in brackets): nos. 1, 3 to 8, 11 to 13, 22, 24, 26 (Aldrich, Steinheim, Germany), no. 2 (Fluka, Neu-Ulm, Germany), no. 16 (Haarmann & Reimer, Holzminden, Germany), nos. 28 and 29 (Lancaster, Eastgate, UK). The following reference substances were synthesized according to the literature: 3-methyl-2-butenthiol (6),

<sup>\*</sup>To whom correspondence should be addressed.

Table 1 GCO-H of roasted coffee powders and brews

No. of compound*	Odour description <sup>b</sup>	RI on RTX-5	FD-factor <sup>c</sup>						
			Po	wder	Brew				
			Arabica	Robusta	Arabica	Robusta	Soluble coffee		
1 Acetaldehyde <sup>d</sup>	Fruity, pungent	<400	25	25	125	62.5	12.5		
2 Methanethiol <sup>d</sup>	Putrid, cabbage-like, sulphurous	<400	5	12.5	1	1	n.d.		
3 Propanal	Fruity	~450	5	5	25	25	25		
4 Methylpropanal <sup>d</sup>	Fruity, malty	<b>~</b> 550	5	5	25	25	25		
5 2,3-Butanedioned	Buttery	580	62.5	125	125	125	62.5		
6 3-Methylbutanal <sup>d</sup>	Malty	653	12.5	25	62.5	62.5	25		
7 2-Methylbutanal <sup>d</sup>	Malty	662	5	12.5	25	25	12.5		
8 2,3-Pentanedioned	Buttery	697	125	125	125	62.5	62.5		
9 3-Methyl 2-butenthiol	Foxy, skunky	822	62.5	62.5	5	5	1		
10 Unknown	Meaty, roasty	839	n.d.	5	n.d.	n.d.	n.d.		
11 2-Methyl-3-furanthiol <sup>e</sup>	Boiled meat-like	870	25	125	5	5	n.d.		
12 Methional <sup>e</sup>	Boiled potato-like	906	62.5	62.5	25	25	5		
13 2-Furfurylthiol <sup>e</sup>	Roasty	911	62.5	125	12.5	12.5	n.d.		
14 Unknown	Roasty	939	n.d.	5	n.d.	n.d.	n.d.		
15 Dimethyltrisulphide <sup>e</sup>	Cabbage-like	970	n.d.	1	12.5	25	n.d.		
16 1-Octen-3-one <sup>e</sup>	Mushroom-like	978	n.d.	n.d.	5	1	n.d.		
17 Unknown	Roasty	986	1	25	n.d.	ī	n.d.		
18 Unknown	Musty, earthy	1002	n.d.	n.d.	5	5	n.d.		
19 Unknown	Earthy, roasty	1012	n.d.	n.d.	5	5	n.d.		
20 3-Mercapto-3-methylbutylformate <sup>c</sup>	Catty, roasty	1022	62.5	62.5	1	1	n.d.		
21 2-Ethyl-3,5-dimethylpyrazine <sup>e</sup>	Earthy, roasty	1086	25	62.5	25	62.5	25		
22 Guaiacol <sup>e</sup>	Phenolic, burnt	1092	12.5	25	25	12.5	5		
23 Unknown	Earthy, roasty	1107	25	125	25	25	5		
24 2,3-Diethyl-5-methylpyrazine <sup>e</sup>	Earthy	1155	25	125	25	125	12.5		
25 Unknown	Earthy	1182	12.5	62.5	1	25	12.5		
26 2-Isobutyl-3-methoxypyrazine <sup>e</sup>	Green, earthy	1186	25	1	125	1	n.d.		
27 Unknown	Roasty, sulphurous	1225	1	5	1	5	n.d.		
28 4-Ethylguaiacol <sup>e</sup>	Phenolic, spicy	1282	n.d.	n.d.	n.d.	5	n.d.		
29 4-Vinylguaiacol <sup>e</sup>	Phenolic, spicy	1317	n.d.	n.d.	5	25	n.d.		
30 (E)-β-Damascenone <sup>e</sup>	Honey-like, fruity	1400	5	5	n.d.	n.d.	n.d.		

<sup>&</sup>lt;sup>a</sup> The compounds which appeared in one of the coffee samples with an FD-factor of at least five are compared.
<sup>b</sup> Odour description assigned during GCO-H.

n.d.: not detectable in the highest headspace volume of 25mL (Table 2).

dimethyltrisulphide (7), 3-mercapto-3-methylbutylformate (2), 2-ethyl-3,5-dimethylpyrazine (8), (E)-B-damascenone(9).

### Gas chromatography-olfactometry

GCO-H was performed with a CP-9001 gas chromatograph connected to the purge and trap system TCT/PTI 4001 (Chrompack, Frankfurt, Germany). The TCT/PTI 4001 system was programmed and controlled via the keyboard of the gas chromatograph. The empty glass tube in the desorption heating block of the purge and trap facility was deactivated by treatment with a mixture of 1,3-diphenyl-1,1,3,3-tetramethyldisilazane, hexamethyldisilazane and pentane (1:1:2, vlwV) (H. Guth, personal communication). The gas chromatograph was equipped with a cooling system for the oven (Chrompack) and with a RTX-5 (crosslinked SE-54)

fused silica capillary (30 m  $\times$  0.52 mm, film thickness 1.5 µm, Amchro, Sulzbach/Taunus, Germany). At 1.6 exit of the capillary, the effluent was split (1 + 1, v/v) into a FID and a sniffing port by using deactivated fused silica capillaries (30 cm  $\times$  0.10 mm). The FID and the sniffing port were held at a temperature of 200 °C. Nitrogen (20 mL/min) was used as make-up gas for the FID. After each run the purge system was automatically cleaned (clean-up flow: 50 mL helium clean-up temperature: 275 °C).

Coffee powder (100 mg) was put into a vessel (volume: 250 mL), sealed with a septum and then held at room temperature for 30 min. Coffee brew (10 mL) was put into a vessel (volume: 100 mL), sealed with a septum and then held in a water bath at 70 °C for 30 min.

The headspace volumes given in Table 2 were drawn by a gas-tight syringe and then injected with a velocity of about 20 mL/min into the purge system which operated in the desorption mode for 10 min at a temperature of

The relationship between FD-factor and headspace volume is given in Table 2.

<sup>&</sup>lt;sup>6</sup> The compound was identified by comparing it with the reference substance using the following criteria: retention index (RI) on the capillary RTX-5, the mass spectrum and the odour quality perceived at the snifting port.
<sup>7</sup> The compound was identified by comparing it with the reference substance on the basis of the RI on capillary RTX-5 and

The compound was identified by comparing it with the reference substance on the basis of the RI on capillary RTX-5 and the odour quality perceived during GCO-H.

Table 2 Injected headspaces volumes and corresponding FD-factors

Headspace volume (mL)	25	5	2	1	0.4	0.2
FD-factor	1	5	12.5	25	62.5	125

280 °C. The carrier gas helium (flow: 20 mL/min) swept the headspace sample into the trap (20 cm × 0.53 mm fused silica capillary coated with CP-sil 8CP, film thickness 5 µm which was cooled with liquid nitrogen at –110 °C. To start the GC run, the trap was heated up very rapidly to 250 °C. This temperature was held for min, and the sample was flushed by the helium (flow rate: 8 mL/min) onto the RTX-5 capillary. The temperature of the oven was held at 0 °C for 2 min, then raised at a rate of 6 °C/min to 50 °C. held for 2 min and then raised at a rate of 8 °C/min to 50 °C. held for 2 min and

### Headspace - GC/MS

The apparatus used for GCO-H was modified. The RTX-5 capillary was replaced by a DB-5 fused silica capillary (30 m × 0.32 mm, film thickness 0.25 μm; 1.8 W. Scientific, Polsom, U.S.A.). The exit of the capillary was coupled with the mass spectrometer Incos XL (Finnigan, Bremen, Germany). The flow of the carrier gas helium was 2 mL/min, and the same temperature program as reported under Headspace-GC/O was used. Headspace sampless of 20 mL were analysed as reported above. Mass spectra were generated in the electron impact mode (MS/EI) at 70 c.

### Results and Discussion

The results of GCO-H of the coffee samples are summarized in Table 1. Twenty odorants (nos. 1 to 9, 11 to 13, 20 to 26, 30) showing FD-factors of 5 and higher were perceived in the headspace of Arabica coffee (powder). Odorants nos. 1 to 8 were identified by comparing the RI value, the mass spectrum and the odour quality with the corresponding properties of the reference substance. No mass spectrum was obtained in the cases of nos. 9, 11 to 13, 20 to 22, 24, 26 and 30. These odorants have been identified in the preceding AEDA (1,2) and most of them have been also quantified (10). Consequently, it was easy to identify them during GCO-H by comparing their RI values and their odour qualities with the corresponding properties of the reference substances. Only the earthy, roasty compounds nos. 23 and 25 remained unidentified.

There was a good agreement in the highly volatile odorants with those reported by Holscher and Steinhart (3) for Arabica coffee (peowder). With exception of hydrogen sulphide and dimethylsulphide, all of the odorants (nos. 1, 2, 4 to 9, 12, 13 in Table 1) sniffed by the odorants (nos. 1, 2, 4 to 9, 12, 13 in Table 1) sniffed by the odorants (nos. 1, 2, 4 to 9, 12, 13 in Table 1) sniffed by the odorants (nos. 1, 2, 4 to 9, 10, 12, 13 in Table 1) sniffed by the odorants (and the odorants) with the odorants (and the odorants)

ing experiments (Guth and Grosch, unpublished results), these odorants are only detectable by GCO-H when the empty glass tube in the desorption block is deactivated (cf. Materials and Methods).

With exception of no. 26, all odorants with an FD-factor of at least five in the headspace of the Arabica coffee occurred also in that of the Robusta coffee (powder), but the latter contained three further odorants (nos. 10, 14, 17) of unknown chemical structure.

On the basis of their high FD-factors, 2,3-butanedione, 2,3-pentanedione, 3-methyl-2-butenthiol, methional, 2-furfurylthiol and 3-mercapto-3-methylbutylformate were the key, highly volatile odorants of both powders. The FD-factors of these and of the odorants nos. 1 to 4. 6, 7, 21, 22 and 30 were identical in both coffee samples. as they differed, at the most, by only one dilution step. which is within the limit of error of the GCO-H. Only quantitative measurements as performed in (10) could make clear whether the concentration levels of these odorants are different in the coffee samples. In contrast, clear differences in the odour potencies were found for 2-methyl-3-furanthiol, 2,3-diethyl-5-methylpyrazine and the unknown no. 23, which all belong only to the key odorants of the Robusta coffee powder on the basis of their high FD-factors (Table 1).

Brews were prepared from both coffee powders and held at 70°C until the headspace samples were drawn and analysed by GCO-H. This was performed very carefully, as the high water pressure at 70°C might damage the stationary phases of the GC capillaries. The results listed in Table 1 indicate that the powders and the brews agreed in compounds nos. 5 and 8 as major odorants. The well-known, strong difference between the overall odours of the powders and the brews was mainly due to a decrease of the FD-factors of the thiols nos. 2. 9, 11, 13 and 20 as well as to an increase in the FD-factors of acetaldehyde, propanal, methylpropanal, 3-methylbutanal and dimethyltrisulphide in the brews. Furthermore, the guaiacols 28 and 29 were detectable in the brew of Robusta coffee, and 29 also in that of the Arabica coffee

A brew prepared from a soluble coffee powder was also analysed by GCO-H. The results in Table 1 reveal that the potency of the major odorants nos. 3 to 8, 21, 22, 24 and 25 agreed with that of the corresponding odorants of either the Arabica or the Robusta coffee brew or of both. On the other hand, the brew prepared from the soluble coffee powder was found to lack acetaldehyde, the thiols nos. 9, 11, 13, methional, unknown no. 23 and the guaiacols 28 and 29 compared to the brews prepared from the Arabica and the Robusta coffee samples.

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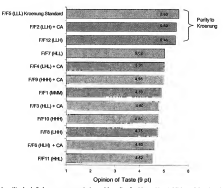
# **APPENDIX F**

### 3.2 Enhanced Roast Coffee (outside standard of identity)

In this case, I I prototypes were produced by adding combinations of 3 chemicals (linalool, 8damaseanone and phenyelthanol) into the already brewed coffee following an experimental design. Base used for the addition of chemicals was standard Jacobs Kroenung that was used as reference as well.

When results are analysed for all consumers, only 2 products scored parity with Kroenung although numerically inferior (Chart 6). All the rest scored significantly lower for this category of consumers.

Chart 6: Overall liking for all consumers



Products codes: (linalool, B-damascanone and phenyelthanol) L= None, H = Addition of the chemical

Chart 7 reflects the results when the analysis is focused in fruity/floral likers (40%, identified through the classification set referred previously). In this case, 4 products scored numerically better than the reference, with one of them being significantly superior. This superiority is driven by the presence of linalood in the Kroenung base and it can be seen in most products with the high scores. The prototypes with linalool scoring lower tend to be those having additional acidity (citric acid). This confirms the sensory assessment of the acidity-fruitiness axis explained in the previous section where fruity/floral likers are described as fruity likers and acidity disilkers. On the other side, Chart 8 shows the tolerance of the fruity/floral disilkers to acidity and their disilke for fruitiness with products with linalool scoring low and products with added acidity spreaded across the scores.

Chart 7: Overall liking for fruity/floral likers (40% all consumers)

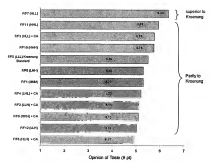
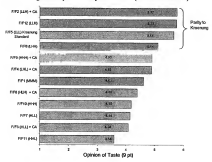


Chart 8: Overall liking for fruity/floral acceptors/dislikers (60% all consumers)



# **APPENDIX G**

### Linalool

From Wikipedia, the free encyclopedia

Linalool (IPA: /li'næloopl/) is a naturally-occurring terpene alcohol chemical found in many flowers and spice plants with many commercial applications, the majority of which are based on its pleasant scent (floral, with a touch of spiciness). It has other names such as β-linalool, linalyl alcohol, linaloyl oxide, p-linalool, allo-ocimenol and 2,6-dimethyl-2,7-octadien-6-ol.

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- 2 Enantiomers
- 3 Biosynthesis
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### In nature

Over 200 species of plants produce linalool, mainly from the families Lamiaceae (mints, scented herbs), Lauraceae (laurels, cinnamon, rosewood) and Rutaceae (citrus fruits), but also birch trees and other plants, from tropical to boreal climate zones. It was also found in some fungi.

Lin	alool							
HQ /								
IUPAC name	[show]							
Identifiers								
CAS number	78-70-6							
PubChem	6549							
SMILES	[show]							
InChI	[show]							
Prop	erties							
Molecular formula	C <sub>10</sub> H <sub>18</sub> O							
Molar mass	154.25 g/mol							
Density	0.858 - 0.868 g/cm <sup>3</sup>							
Melting point	< 20 °C							
Boiling point	198 – 199°C							
Solubility in water	1.589 g/l							
Haz	ards							
Flash point	55 °C							
materials in the	erwise, data are given for ir standard state , 100 kPa) references							

### Enantiomers

Linalool has a chiral center at C<sub>3</sub> and therefore two stereoisomers: **licareol** is (R)-(+)-linalool with CAS No. 126–90–9 (PubChem 67179) and **coriandrol** is (S)-(-)-linalool with CAS No. 126–91–0 (PubChem 13562).

Both enantiomeric forms are found in nature: S-linalool is found, for example, as a major constituent of the essential oils of coriander (Coriandrum sativum L. family Apiaceae) seed, palmarosa [Cymbopogon martinii (Roxb.) Wats, family Poaceae], and sweet orange (Citrus sinensis Osbeck, family Rutaceae) flowers. R-linalool is present in lavender (Lavandula officinalis Chaix, family Lamiaceae), laurel (Laurus nobilis, family Lauraceae), and sweet basil (Ocimum basilicum, family Lamiaceae), among others

Each enantiomer evokes different neural responses in humans, and therefore are anthropophilically classified as possessing distinct scents. 3S-(+)-linalool is perceived as sweet, floral, petitgrain-like (odour threshold 7.4 ppb) and the 3R-form as more woody and lavender-like (odour threshold 0.8 ppb)

### Biosynthesis

In higher plants linalool as other monoterpenoids is produced from isopentenyl pyrophosphate via the universal isoprenoid intermediate geranyl pyrophosphate, through a class of membrane-bound enzymes named monoterpene synthases. One of these, linalool synthase (LIS), has been reported to produce (S)-linalool in several floral tissues.

### Uses

In addition to its use as a scent in domestic products such as soap, detergent, shampoo, and lotion, linalool is also used as a chemical intermediate. One common downstream product of linalool is Vitamin E. Linalool, is also used by pest professionals as a flea, and cockroach insecticide

### Safety information

Linalool should be avoided by people with perfume allergy[1].

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### Notes

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### External links

- Comprehensive data sheet (http://www.inchem.org/documents/sids/sids/78706.pdf)
- Record (http://householdproducts.nlm.nih.gov/cgi-bin/household/brands?tbl=chem&id=2771) in the Household Products Database of NLM

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